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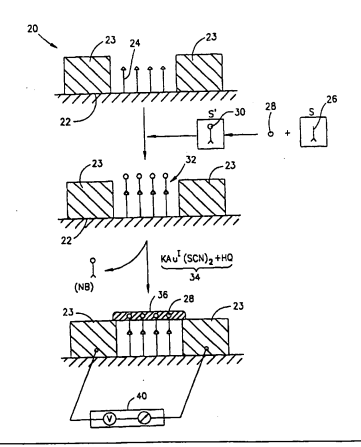
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(54) Title: METHOD FOR GOLD DEPOSITION

(57) Abstract

A method for depositing gold on a substrate is provided. By this aspect, nucleation centers are bound, deposited or formed at one or more sites on the substrate, the substrate is then contacted with a treatment composition which comprises soluble gold-providing agent from which gold is deposited onto a nucleation center to yield gold metal deposits. The method of the invention may be used for a variety of image enhancing techniques as well as diagnostic and analyte detection techniques.



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METHOD FOR GOLD DEPOSITION

FIELD OF THE INVENTION

The invention relates to chemical deposition of gold metal on substrates.

BACKGROUND OF THE INVENTION

There are various techniques and uses in which metal substances are deposited on substrates. In various such methods, the metal to be deposited is provided in a form of a soluble complex or molecule, e.g. in an aqueous solution. Under appropriate conditions the complex or molecule is reduced and deposits the metal onto the substrate.

At times, in order to ensure deposition of metal from solution onto a substrate, it is preferable to first provide nucleation centers on the substrate. Such nucleation centers may, for example, be a variety of metal complexes, clusters or small metal particles, deposited on or attached to the surface.

Examples of techniques involving metal deposition are a variety of image formation or latent image contrast-enhancement techniques for photography, optical microscopy or electron microscopy. Another example of laboratory techniques where metal complexes are deposited onto a substrate, is in imaging of separation products, e.g. proteins or nucleic acids, in a variety of chromatographic or electrophoretic techniques.

The metal most typically used in such techniques is silver. However, one typical drawback of silver deposition is that the deposition is not completely specific and some degree of spontaneous deposition of silver may occur also without an *a priori* deposition of nucleation centers. Thus, where silver deposition is used in image-enhancement or contrast-enhancement methods, there is typically a limited signal-to-noise ratio.

GENERAL DESCRIPTION OF THE INVENTION

The present invention has, as one of its objects, the provision of a method for depositing gold onto specific sites on a substrate. Another object of the invention is to provide a composition and kit for use in such a method.

In accordance with the present invention gold, initially contained in a soluble gold-providing agent, being a gold-containing molecule, cluster or complex, is deposited onto nucleation centers. The term "nucleation centers" as used herein refers to any salt, complex, cluster or particle having catalytic properties of transforming gold from a gold-containing molecule or complex into gold metal. Particles with a metal surface are, for example, colloid metal particles, a variety of metal complexes, clusters containing metal atoms, etc.

In accordance with the invention there is provided a method for depositing gold at one or more sites on a substrate, comprising:

- binding, depositing or forming nucleation centers at said one or more (a) sites;
- contacting said one or more sites with a treatment composition (b) comprising said gold-providing agent and a reagent, the composition being kinetically stable such that gold metal is essentially not deposited on the substrate unless a nucleation center is present on the substrate and in the presence of a 20 nucleation center at said one or more sites, gold atoms are released from said gold-providing agent and deposited onto said nucleation center to form gold metal deposits at said one or more sites.

As will be appreciated, once gold is deposited on the substrate, it serves as a nucleation center for further gold deposition. By appropriately controlling the 25 incubation time as well as other conditions affecting the gold deposition (some of which will be outlined bellow), the amount of the deposited gold at a deposition site can be controlled. Furthermore, although the initial gold deposition may be at discrete sites on the substrate, by allowing the gold deposits to grow, following further deposition, discrete sites may combine to form one larger gold deposit 30 encompassing the so combined sites.

The term "essentially not deposited" means to denote that there is either no gold separation from the solution and the gold atoms remain within the soluble molecule or complex, or that gold metal deposition on a substrate is negligible and thus not or only hardly observable and measurable, yielding a high signal-to-noise ratio.

In accordance with one embodiment, the nucleation centers are formed by chemical or physical deposition of nucleation center forming agent onto the one or more sites. By another embodiment, such nucleation centers may be formed chemically at said one or more sites by a redox reaction in which metal ions, in particular silver ions, are oxidized into metallic silver. Such chemical formation of nucleation centers is known *per se* (see for example WO 99/04402 and PCT/IL99/00232).

In accordance with yet another embodiment, the nucleation centers are formed by binding of nucleation center-forming agents at said one or more sites. Such agents comprise at least one binding moiety with a binding affinity (which may be specific or non-specific) to said one or more sites, which is bound to at least one nucleation center moiety. The nucleation center moiety is a species that can act as a catalytic agent for gold metal deposition, e.g. a salt, metal particle, cluster of metal atoms or a metal-containing complex. The binding moiety may, for example, be one member of a recognition group, where another member of the recognition group forms or is included in said one or more sites. The recognition group, typically a recognition couple, consists of two or more substances with a binding affinity one to the other. The recognition group may be, but is not limited to, any one of the following couples: an antigen and an antibody or an antibody derivative 25 with a complementary antigen-binding domain; sugar and a lectin; a receptor and a ligand; a nucleotide sequence and a complementary nucleotide sequence; a nucleotide sequence and its binding protein or synthetic binding agent; biotin and avidin or streptavidin; cellulose or chitin and cellulose binding domain.

The nucleation center moiety is preferably a gold particle or a cluster containing gold atoms, e.g. Au₁₁ or Au₅₅ or Au₁₄₇ (gold clusters that contain 11, 55

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or 147 gold atoms, respectively). The treatment composition is typically an aqueous solution. Said gold-providing agent is preferably a gold complex such as Au^l(SCN)₂ usually in the form of an alkali metal (e.g. Na⁺ or K⁺) or ammonium salt.

The reagent in the treatment composition may be a reducing agent. Where said gold-providing agent is Au¹(SCN)₂ the reagent is preferably a quinone, e.g. hydroquinone (HQ) or naphtohydroquinone (NHQ). However, any other agent which can reduce a gold-containing molecule or complex to obtain gold metal, but because of kinetic limitations (e.g. high activation energy and low pre-exponential factor) will do so in a negligible rate in the absence of a nucleation center, yet, in a meaningful rate (namely in a rate that yields observable results in a not too long period of time) when a nucleation center is present, may also be used.

A treatment composition comprising Au^I(SCN)₂ and HQ in an acidic medium is stable, meaning that gold remains as soluble species in solution and neither precipitates as solid metal nor is deposited on solid substrates other than nucleation centers. When the pH of such a treatment composition is below about 6, the treatment composition is particularly stable. Upon contact with a nucleation center gold metal is deposited on sites of the substrate comprising nucleation centers.

The rate of deposition of the gold metal may be controlled, for example, by controlling the pH of the treatment composition: increase in pH will increase the rate of gold deposition and a decrease in pH will decrease the deposition rate. Furthermore, the rate of deposition may also be controlled by a variety of other means, for example: by controlling the flow parameters of the treatment 25 composition (rapid flow prevents poisoning by by-products, such as CN, which can reduce the rate of gold deposition); by adding surface active agents which can either increase or decrease the rate of deposition (for example polyamines, poly acids or polyalcohols and other surface active agents may have the effect of altering the rate of gold deposition). In addition, by the use of various additives, e.g. those noted above, the roughness of the surface of the deposited gold may be changed

(such a change may result in a change of color and electrical properties of the deposited gold).

The method of the invention may be used for assaying the presence or concentration of a specific substance at sites on a substrate. Such a method comprises the following steps:

- (a) providing conditions allowing formation of nucleation centers on sites containing said substance;
- (b) contacting said substrate with a treatment composition comprising said gold-providing agent and a reagent, the composition being kinetically stable such that upon such exposure gold metal is essentially not deposited on the substrate unless a nucleation center is present thereon, and in the presence of a nucleation center at said sites, gold atoms are released from said gold-providing agent and deposited onto said nucleation centers to form gold metal at said sites; and
 - (c) detecting gold deposits on said substrate, a gold deposit at a site on the substrate indicating presence of said substance at said site.

In accordance with another embodiment, the gold deposition method of the invention is used in an assay intended to detect the presence of an analyte in a sample. In particular, the present invention is applicable to such a method where a capturing agent held on a substrate is used to detect the presence of an analyte in a

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sample that binds specifically to the capturing agent. Such a capturing agent may be a member of a recognition group, while the analyte is another member of the group. A specific example is the case of assaying for the presence of a target oligonucleotide in a sample, with the capturing agent being an oligonucleotide with a sequence complementary thereto. A nucleation center forming moiety may be bound to analytes presence in the sample, e.g. to the oligonucleotides in the sample, and then after allowing the sample to react with the substrate, and performing a gold deposition reaction as described above, formation of gold deposits will indicate the presence of the analyte in the sample.

A method of the invention of preparing a specimen for imaging in a microscope, comprises the following steps:

- (a) providing conditions permitting formation of nucleation centers at selective sites of the specimen; and
- (b) contacting said specimen with a treatment composition comprising said gold-providing agent and a reagent, the composition being kinetically stable such that gold is essentially not deposited on the specimen unless a nucleation center is present thereon, whereby gold atoms from said gold-providing agent are released and deposited onto said specimen at said selective sites.

A method of the invention for detecting the presence of an analyte in a sample, comprises the following steps:

- (a) treating the sample to bind nucleation center forming moieties to the analyte, if present in the sample;
- (b) reacting the sample with a substrate, holding capturing agents that specifically bind said analyte, and removing unbound sample portions;
- (c) contacting said substrate with a treatment composition comprising said gold-providing agent and a reagent, the composition being kinetically stable such that upon such exposure gold metal is essentially not deposited on the substrate unless a nucleation center is present thereon, and in the presence of a nucleation center at said sites, gold atoms are released from said gold-providing

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agent and deposited onto said nucleation centers to form gold metal at said sites; and

(d) detecting gold deposits on said substrate, a gold deposit indicating presence of the analyte in the sample.

A specific application of this method is in the oligonucleotide chips. Oligonucleotide chips have different capturing oligonucleotides deposited in different sites on the chip. Applying the above method may allow to simultaneously assay for a large number of different oligonucleotides in a sample.

In a so prepared specimen, said selective sites thus contain gold deposits which permit their visualization. Said selective sites may typically be sites containing a specific substance and the nucleation centers are then formed at said sites by nucleation center forming agents with a moiety having specific binding affinity to said specific substance. In this manner, for example, specific cells, organelles, regions rich with a specific substance, etc. can be clearly visualized.

A method according to the invention for identifying locations of a specific separation product on a substrate, comprises the following steps:

- (a) providing conditions permitting formation of nucleation centers at sites on said substrate comprising said specific separation product;
- (b) contacting said substrate with a treatment composition comprising said gold-providing agent and a reagent, the composition being kinetically stable such that gold is deposited on the substrate essentially only where a nucleation center is present thereon, whereby upon such contact, gold atoms from said gold-providing agent are released and deposited onto said nucleation center to form gold metal; and
 - (c) detecting presence of gold on said substrate, such gold signifying presence of said specific separation product on the substrate at a site where the gold is detected.

In a separation technique where the sample is separated into various fractions based on different movement rates of different substances through a medium (typically a gel or a solid) under applied conditions, for example

electrophoresis, thin layer chromatography (TLC), etc., the nucleation centers-forming agents may be admixed to the sample prior to separation. This is usually preferred. For one such prior addition of the nucleation center-forming agent may be more economical, as less material may be required. In addition, where such agents are applied to the substrate after the separation, they have to diffuse through the medium in order to reach the separated substance located within the medium; such diffusion may be a limiting factor in proper gold-staining of separated fractions. However as will readily be appreciated, it is possible at times to add the nucleation center-forming agent also after separation.

The present invention also provides a method for assaying presence of an analyte in a sample. Such a method comprises the following steps:

- (a) providing a substrate carrying capturing agents which bind said analyte;
- (b) contacting said substrate with said sample, whereby said analyte binds to said capturing agents;
 - (c) providing conditions permitting formation of nucleation centers on sites of said substrate comprising said analyte;
- (d) contacting said substrate with a treatment composition comprising said gold-providing agent and a reagent, the composition being kinetically stable such that gold is deposited on the substrate essentially only at sites thereof containing the nucleation centers; and
 - (e) detecting metallic gold deposits on said substrate, indicative of the presence of said analyte in said sample.

The detection in step (e) of the method for assaying presence of an analyte in a sample, may be carried out by testing appearance of gold deposited on the substrate (gold deposited, depending on size and roughness, may have a color ranging from yellow to orange to red).

In accordance with one embodiment of the invention, the detection in step (e) is based on the electric conductivity of the deposited gold. In this embodiment the capturing agents may be carried on the substrate between

electrodes or may be carried on one or more electrodes such that the gold deposited onto the nucleation center in step (d) establishes an electric contact between electrodes. The detection of the gold deposits in step (e) is thus carried out by determining existence of electrical contact between the electrodes, e.g. by measuring the current-potential relationship between the electrodes. The magnitude of the current-potential relationship may be used as a gauge for the agent's concentration in the sample. Alternatively, determining the number of connections in a large array of similar electrode couples with identical capturing agents carried between or on them, may also provide a quantitative measure of the analyte concentration.

The assayed analyte may be a member of any of the recognition couples mentioned above, with the other member of the couple being then included as a moiety in the capturing agent. The formation of the nucleation centers in step (c) may be the provision of agents having a binding moiety with specific binding affinity to the analyte captured on the substrate by the capturing agent, bound to a nucleation-forming moiety, which may be a metal particle, a cluster containing metal atoms or a metal-containing complex, or any other species having the catalytic properties of transforming gold-containing molecular species into gold metal. For example, where the analyte is one member of a binding couple, the 20 binding moiety may be another member (e.g. where the analyte is an antigen, the nucleation center- forming agent may comprise an antibody, which may be the same or different as an antibody used as the capturing agent). The present invention further provides a kit for use in the above method. Such a kit typically comprises a treatment composition. In addition, such a kit may also comprise a reagent for forming nucleation centers at specific sites on a substrate, e.g. such having a specific binding moiety being a member of any of the above binding couples. In addition, a kit for use in the above assay method may also comprise, according to some embodiments, a reagent system for tagging an analyte in a sample with nucleation center-forming moiety. Furthermore, in accordance with some

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embodiments of the invention, the kit may comprise a substrate comprising electrodes with capturing agents disposed between and/or on the electrodes.

In order to understand the invention and to see how it may be carried out in practice, a preferred embodiment will now be described, by way of non-limiting example only, with occasional reference to the accompanying drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

- Fig. 1 shows reaction schemes of the manner of preparation of a treatment composition in accordance with an embodiment of the invention (Scheme (a) and Scheme (b)) and of gold deposition onto a nucleation center (Scheme (c)).
- Fig. 2 is a schematic representation of an assay of an analyte in a sample in accordance with an embodiment of the invention.
- Fig. 3 is a schematic representation of an assay in accordance with another embodiment of the invention.
- Fig. 4 is a schematic representation of an assay in accordance with a further embodiment of the invention.
 - Fig. 5 is a schematic illustration of the manner of enhancing specific bands on a solid or a gel substrate, separated by chromatography or by electrophoresis.
 - Fig. 6 is a schematic illustration of a method in accordance with an embodiment of the invention for imaging specific parts of a microscope specimen.
- Fig. 7 shows the application of the system in the detection of analytes, oligonucleotides with a specific sequence in this specific example, in a sample.

DETAILED DESCRIPTION OF A PREFERRED EMBODIMENT Preparation of a treatment composition and general gold deposition scheme

A treatment composition in accordance with the preferred embodiment of the invention comprises a gold-containing complex- Au^ISCN. In order to prepare such a treatment composition at a first stage a solution comprising KAu^{III}Cl₄ and KSCN is prepared, typically by mixing two stock solutions, one comprising the

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gold containing complex and the other the KSCN. For example, such a solution may be formed by mixing equal volumes of a first solution comprising about 0.5 M KSCN and a second solution comprising about 0.05 M KAu^{III}Cl₄. Such mixing results in the formation of the complex KAu^{III}(SCN)₄ (See Scheme (a) in Fig. 1). This complex precipitates but spontaneously forms the complex KAu^I(SCN)₂. Typically, after the KAu^{III}(SCN)₄ is precipitated, it is washed and resuspended in water or buffer and allowed to undergo the spontaneous reaction (SPONT.) noted above (illustrated in Scheme (b) of Fig. 1).

The KAu^I(SCN)₂ solution is then mixed with a reducing agent, e.g. with about 0.055 M hydroquinone (HQ). The reduction of the gold-containing complex by the hydroquinone is then thermodynamically favorable, but owing to the large activation energy and low pre-exponential factor, the kinetics is very slow. At a pH of less than about 6, the spontaneous reduction is practically nil. Thus, the solution is stable and gold metal is not deposited spontaneously.

However, as shown in Fig. 1, scheme (c), upon exposure to nucleation centers (NC), acting as catalysts, which may for example be gold colloids, other metal colloids, clusters of metal atoms, metal-containing complexes, metal ions, etc., gold metal (Au⁰) is being deposited on the surface of the nucleation center associated with the formation of benzoquinone (BQ).

The pH level controls the rate of gold deposition: acidification of the solution, namely lowering pH, slows the metal deposition, while addition of a base, namely increasing pH, accelerates the rate of metal deposition.

It should be noted that under basic pH, some spontaneous deposition of gold may take place even without nucleation centers.

Various additives which may either be added initially to the treatment solution, or added during the deposition process, may influence the rate of the gold deposition process and the pattern of the gold deposit (size of deposits, surface roughness, color, etc.). For example, addition of halogen ions to the treatment composition results in surface poisoning of the growing metal centers. This will reduce the rate, and eventually stop the gold deposition process. Use of

such additives thus allows a control of the deposited gold crystalinity, grain size, the rate of the gold deposition, the roughness of the surface of the deposited gold, the electrical properties, etc. The gold deposition process may obviously also be controlled by the change of various other parameters such as temperature, oxygen content, light flux, solution-agitation rate, etc.

The gold deposition process of the invention can be used for a variety of diverse applications, some examples of which will be illustrated below.

Detection of analytes in a sample

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An assay in accordance with one embodiment of the invention is illustrated in Fig. 2. Provided by this embodiment is a device 20 having a non conductive substrate 22 carrying electrically conductive electrodes 23. In the gap between electrodes the substrate 22 carries capturing agents 24 which have a specific binding affinity to analyte 26 to be assayed.

Sample S, which comprises analyte 26, is pretreated by adding nucleation centers 28 and treating it in a manner so that nucleation centers will become coupled to the analyte in a covalent or non covalent interaction, to yield modified analyte 30. In the case of an oligonucleotide analyte, such coupling may, for example, be performed by heating the sample in the presence of a 20 cis-platinum-biotin and then adding an appropriately derivatized cluster, e.g. Au₅₅(Ph₃P)₁₂Cl₆-streptavidin cluster. The cis-platinum-biotin binds to the oligonucleotides, then Au₅₅(Ph₃P)₁₂Cl₆- streptavidin binds to the biotin moieties on said nucleotides, thus binding nucleation moieties to the analyte. In the case of a non-nucleotide analyte, coupling may be achieved, for example by binding 25 amino-derivatized cluster or colloid to carboxy moieties of the analyte; coupling carboxy-derivatized clusters or colloids to amino-moieties of the analyte; coupling maleimido-derivatized cluster or colloids to thiol moieties of the analyte; etc. As will be appreciated, other substances in a sample may also become tagged with the nucleation center in this process. However, these will not bind to capturing agents 24 and the nucleation centers coupled thereto will thus not remain on the substrate after treatment. (See description below).

Upon contact with device 20, modified analyte 30 binds to the capturing agents to yield nucleation center-forming complexes 32 which are immobilized on substrate 22. Washing the substrate removes unbound substances (NB).

Device 20 may then be contacted with the treatment solution 34 which comprises a gold containing complex such as KAu¹(SCN)₂ and a reducing agent such as hydroquinone (HQ) and consequently upon contact with the nucleation centers 28, gold is deposited on the nucleation centers and forms a continuous gold mass 36 extending between electrodes 23. In order to avoid that electrodes 23 will act as nucleation centers, they may either be made of a conductive material lacking catalytic properties, e.g. silicon passivated with silica, with long alkyl chains, or passivated by coating with an inert material, or, where the electrodes are made of a metal, they may be pretreated to render them inert as gold-deposition catalysts, e.g. by incubation with a solution comprising a long chain alkane thiol, such as 1-octadecyl-thiol.

By measuring current voltage relationships using a measurement device 40, the existence of an electric contact between the electrodes 23 can be determined, the presence of which indicates the presence of the analyte in the assayed sample.

As can generally be appreciated by the artisan, the binding of the capturing agents to the substrate may be achieved in a number of ways. For example, oxide surfaces may be derivatized with a silicon reagent such as (CH₃CH₂O)₃Si-R-X, wherein R may be an alkyl, aryl or other spacer and X may be an active moiety for subsequent binding to a moiety Y in the capturing agent. The nature of X and Y depends on the couple and may be selected from couples consisting of acid and amine, hydroxy and acid, cycloaddition reactions, radical reactions, nucleophilic substitutions, non- covalent interactions, etc., as is well known and clear to the artisan. Where the substance is made of a polymeric material, binding, by any of

the above type of reactions, or others, may also be achieved between the capturing agent and a side group of the polymeric material.

Device 20 may comprise two or more electrodes, typically a plurality of electrodes arranged in an array. The array may assume different geometries. The same capturing agents may be carried between different pairs of electrodes, and/or different capturing agents may be deposited between different electrodes. A device of the latter kind may be used in the multiplexing assay for simultaneous determination of a number of analytes. In addition, where the same capturing agents are carried between different pairs of electrodes, this may allow to perform a quantitative assay of concentration of the analyte: for example, the concentration may be assessed based on the number of electric contacts formed between different pairs of electrodes - a large number of electric contacts signifying a large concentration and a small number corresponding to a small concentration.

The electrode array may also be part of a more complex electronic setup containing different components, e.g. diodes, transistors, conductors, capacitors, etc.

Another embodiment of an assay in accordance with the invention can be seen in Fig. 3. The main difference in the case of the embodiment of Fig. 3 from that described in Fig. 2, is that rather than treating sample S to bind nucleation centers 28 to analytes 26 to obtain a modified analyte 30, in the case of the embodiment of Fig. 3, a nucleation center-forming agent 150 is provided. Other components of this assay are very similar to those of Fig. 2 and thus the same reference numeral shifted by one hundred (namely with a "1" prefix) to those used in Fig. 2 were employed in Fig. 3 to indicate like components.

After contact of a device 120 with a sample, analyte 126 binds to a capturing agent 124 and then after washing free (non bound (NB)) analyte molecules and other substances in the sample, the device is contacted with a nucleation center-forming agent 150 which comprises a moiety 152 with a specific binding affinity to analyte 126 which is now bound to capturing

agents 124, coupled to a nucleation center 154. Thus, an immobilized nucleation center 132 forms on the substrate in the gap between electrodes 123. Then, after removal of non bound agents 150 (NB) and adding a treatment composition 134 the gold mass 136 forms which provides electric contact between electrodes 123, which may then be determined similarly as shown in Fig. 2.

A further embodiment of an assay in accordance with the invention is seen Fig. 4. In Fig. 4, like elements to those of Figs. 2 and 3 have been marked with the same reference numeral to that used in Fig. 2 shifted by one hundred (namely with a "2" prefix). Sample S containing analyte 226 is reacted with biotin 227 to yield a treated sample S' with analyte-biotin complexes 231. These are then contacted with substrate 222 which contain capturing agents 224 fitted between electrodes 223, to yield modified capturing complexes 233 on substrate 222.

Gold colloid particles 240 are reacted with an avidin or streptavidin 242 to yield nucleation center forming agents 243. Contacting agent 243 with modified substrate 222 then yields immobilized nucleation centers 235. The subject can then be contacted with a treatment solution 234 similarly as in Figs. 2 and 3 resulting in gold deposition to form a continuous gold mass 236 between electrodes 223.

As will no doubt be appreciated by the artisan, various modifications of the embodiments shown above are possible.

Visualization of separation products

The method of the invention can be used for visualizing separation product obtained in a number of separation techniques such as gel electrophoresis, gel permeation techniques, size exclusion chromatography, affinity chromatography and others. This is illustrated schematically in Fig. 5.

Substrate 200, which may be a gel or a solid substrate containing different lanes 202, each consisting of a plurality of bands 204, each one being a specific fraction of a sample separated in each lane. Such a substrate 200 may be obtained by any of the techniques noted above.

In order to visualize the existence of a specific substance and locate its band, a nucleation center-forming agent 206 which has a moiety 208 which specifically binds to the substance in a nucleation center 210 is contacted with substrate 200 and after access, agents 206 are washed away, the substrate is contacted with a treatment solution 212 and consequently gold deposits 214 will form on bands containing that substance (the other bands are not visible and are included only for illustration purposes). Preferably, however, nucleation center-forming agents 206 may be added to the substrate prior to the beginning of the separation process since diffusion of subsequently-added nucleation center-forming agents may be a limiting factor to achieve binding of these agents to the separated substance.

Depending on the exact deposition conditions, which will *inter alia* determine the roughness of the surfaces of the deposited gold mass, the color of the bands may vary between yellow to black. Using various visualization techniques, based on measurement of light absorbance or transmission through the bands, a quantitative measurement of a separate substance may be obtained.

In the alternative, rather than optical techniques, the gold deposits may also be detected by various electric detection techniques. This may be achieved for example by providing two planar arrays of electrodes sandwiching the separation matrix which will permit an automatic electric detection.

Contrast enhancing in imaging techniques

There are many imaging techniques which request contrasting agents in order to view certain components. Such is the case in optical microscopy, electron microscopy (transmission electron microscopy, scanning electron microscopy) as well as in other imaging techniques, such as atomic force microscopy. Hitherto, such imaging techniques involved silver metal deposition or carbon deposition and some others.

In the case of biological specimens, contrast enhancers are required to allow visualization of specific tissues, cells, organelles, etc.

In accordance with the invention, as illustrated in Fig. 6, a biological specimen 300, containing a cell 302 with a non visible organelle 304, is treated in accordance with the invention by successive exposure to nucleation center-forming agent 306 which has a binding moiety 308 specific to substance 5 included in organelle 304 and a nucleation center 310 and after treatment with a treatment composition 312, organelle 304 is visible as a dark or opaque deposit 314.

Imaging of oligonucleotide hybridization

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The assaying of the presence and concentration of specific-sequence oligonucleotides is nowadays typically being performed by the use of oligonucleotide probes of a complementary sequence attached to known locations on a substrate. Specific and widely used examples of such systems are the so-called DNA-chips. Such chips are surfaces bearing a plurality of oligonicleotide-probe sequences in known and discreet locations on a surface. A sample that may or may not contain the target sequences that complement the probe sequences on chip is treated so that all DNA fragments are labeled (usually by fluorescent labels). Contacting the sample with the surface of the chip under appropriate conditions, the target oligonucleotides attach to the surface in locations bearing the complementary sequence. Detection of hybridization products is usually done by following the fluorescence of labeled duplexes. The location of hybridization product indicates the sequence of the probe and fluorescence intensity indicates their surface concentration.

Reference is now being made to Fig. 7 showing the application of the gold deposition method of the invention in visualization of hybridization products on a DNA-chip using different possible visualization techniques such as optical techniques (absorption, reflectance) and non optical visualization techniques (AMF, STM etc.). As no doubt will be appreciated, this is but an example to the use of the method of the invention for visualizing binding of targets to probes in general, e.g. binding of c-DNA strands to complementary strands, a binding of an antibody to an antigen and other types of recognition couples.

A DNA-chip substrate 400 carries a plurality of DNA probe sites, three 402, 404 and 406 being exemplified, each one having a different nucleotide 5 sequence. A sample S that may contain the target oligonucleotides 410 is treated in a way that all DNA species in solution become labeled by label 412, thus obtaining modified sample S' that may contain modified analyte (if analyte was present in S). Contacting modified sample S' with DNA-chip substrate 400 under appropriate conditions allows selective hybridization of DNA probes on chip to 10 target oligonucleotides if present in modified sample S'. After washing the chip, nucleation centers-forming agents 430 are introduced to the system under condition allowing their binding to labels 412 on labeled oligonucleotides if present at the different sites on chip. After washing, nucleation centers are present only at locations where hybridization between probes on chip and target 15 from solution occurred. Visualization of hybridization is achieved by applying the gold deposition method of the invention, thus growing gold grains from the nucleation centers on-chip and detecting the gold 440 grains using different techniques such as optical techniques and SPM techniques (AFM, STM etc.).

In another embodiment, the label serves itself as a nucleation center. In such case, the step of attaching the nucleation center to the label is skipped.

CLAIMS:

- 1. A method for depositing gold at one or more sites on a substrate, comprising:
- (a) binding, depositing or forming nucleation centers at said one or more sites;
- (b) contacting said one or more sites with a treatment composition comprising a soluble gold-providing agent, being a gold-containing molecule or complex and comprising a reagent, the composition being kinetically stable such that gold is essentially not deposited on the substrate unless a nucleation center is present on the substrate and in the presence of a nucleation center at said one or more sites, gold atoms are released from said gold-providing agent and deposited onto said nucleation center to form gold metal deposits at said one or more sites.
 - 2. A method according to Claim 1, wherein step (a) comprises:
- (a1) providing nucleation center-forming agents which comprise each at
 least one moiety having specific binding affinity to said one or more sites, coupled
 to at least one nucleation center being one or more of the group consisting of metal
 particle, cluster containing metal atoms and a metal-containing complex; and
 - (a2) contacting said one or more sites with said agents.
- 3. A method according to Claim 2, wherein said moiety is a member of a recognition group consisting of two or more molecules or complexes which specifically bind to one another, the other member included in or forming said one or more sites.
- 4. A method according to Claim 3, wherein said recognition group is a member of the group consisting of: an antigen and an antibody or an antibody derivative with an antigen-binding domain; sugar and a lectin; a receptor and a ligand; a nucleotide sequence and a complementary nucleotide sequence; a nucleotide sequence and its binding protein or other specific binding agent; biotin and avidin or streptavidin; cellulose or chitin and cellulose binding domain.

- 5. A method according to any one of Claims 2 to 4, wherein said nucleation center is a gold particle or a cluster containing gold atoms.
 - 6. A method according to any one of Claims 1 to 5, wherein said treatment composition is an aqueous solution.
- 5 7. A method according to Claim 6, wherein said gold-providing agent is $Au^{I}(SCN)_{2}$.
 - 8. A method according to Claim 7, wherein said reagent is hydroquinone or naphtohydroquinone.
- 9. A method for assaying presence of a specific substance at sites on a substrate, comprising:
 - (a) providing conditions allowing formation of nucleation centers on sites containing said substance;
- (b) contacting said substrate with a treatment composition comprising a soluble providing agent, being a gold-containing molecule or complex and comprising a reagent, the composition being kinetically stable such that upon such exposure gold metal is essentially not deposited on the substrate unless a nucleation center is present thereon, and in the presence of a nucleation center at said sites, gold atoms are released from said gold-providing agent and deposited onto said nucleation center to form gold metal deposits at said sites; and
 - (c) detecting metallic gold deposits on said substrate, a gold deposit at a site on the substrate indicating presence of said substance at said site.
 - 10. A method according to Claim 9, wherein step (a) comprises:

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- (a1) providing nucleation center-forming agents which comprise each at least one moiety, having specific binding affinity to said substance, bound to at least one nucleation center being one or more of the group consisting of metal particle, cluster containing metal atoms and a metal-containing complex; and
 - (a2) contacting said substrate with said agents.
- 11. A method according to Claim 10, wherein said moiety is included in one member of a recognition couple consisting of two molecules or complexes which

specifically bind to one another, the other member included in or constituting said substance.

- A method according to Claim 11, wherein said recognition couple is a member of the group consisting of an antigen and an antibody or an antibody derivative with an antigen-binding domain; sugar and a lectin; a receptor and a ligand; a nucleotide sequence and a complementary nucleotide sequence; a nucleotide sequence and its binding protein or other specific binding agent; biotin and avidin or streptavidin; cellulose or chitin and cellulose binding domain; said moiety being one of the recognition couple and said substance being or including 10 the other.
 - A method according to any one of Claims 9-12, for assaying the presence of **13**. one or more target oligonucleotides with a specific target sequence in a sample, comprising:
- providing a substrate carrying one or more probe oligonucleotides (a) each with a probing sequence complementary to one of said target sequences;
 - contacting said substrate with the sample and providing conditions allowing formation of nucleation centers where a target oligonucleotide hybridizes to a probe oligonucleotide;
- contacting said substrate with a treatment composition comprising a gold-providing agent, being a gold-containing molecule or complex and comprising a reagent, the composition being kinetically stable such that upon such exposure gold metal is essentially not deposited on the substrate unless a nucleation center is present thereon, and in the presence of a nucleation center at said sites, gold atoms are released from said gold-providing agent and deposited onto said nucleation 25 center to form gold metal deposits at said sites; and
 - detecting metallic gold deposits on said substrate, a gold deposit at a site on the substrate indicating the presence of said substance at said site.
 - A method according to Claim 13, wherein said step (b) comprises:
 - treating said sample to bind a label to oligonucleotides in the sample; (b1)
- contacting said sample with said substrate; and (b2) 30

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- (b3) contacting said substrate with a nucleation center-containing agent, capable of specific binding to said label.
- 15. A method according to Claim 13, wherein said step (b) comprises:
 - (b1) treating said sample so as to bind nucleation centers to oligonucleotides present therein; and
 - (b2) contacting said samples with said substrate.
- 16. A method according to any one of Claims 9 to 15, wherein said nucleation center is a gold particle or a cluster containing gold atoms.
- 17. A method according to any one of Claims 9 to 16, wherein said treatment composition is an aqueous solution.
 - 18. A method according to Claim 17, wherein said gold-providing agent is $Au^{I}(SCN)_{2}$ and said reagent is a quinone.
 - 19. A method according to any one of Claims 9 to 18, wherein said substrate is a specimen for viewing in an imaging technique.
- 20. A method according to Claim 19, for preparing the specimen for viewing in a microscope, comprising the following steps:
 - (a) providing conditions permitting formation of nucleation centers at selective sites of the specimen; and
- (b) contacting said specimen with a treatment composition comprising a soluble gold-providing agent, being a gold-containing molecule or complex and comprising a reagent, the composition being kinetically stable such that gold is essentially not deposited on the specimen a nucleation center is present thereon, whereby gold atoms from said gold-providing agent are released and deposited onto said specimen at selective sites.
- 25 21. A method according to any one of Claims 9 to 18, wherein said substrate contains separated fractions of a sample.
 - 22. A method according to Claim 21 for identifying locations of a specific separation product on the substrate, comprising the steps of:
- (a) providing conditions permitting formation of nucleation centers at sites on said substrate comprising said specific separation product;

- (b) contacting said substrate to a treatment composition comprising a soluble gold-providing agent, being a gold-containing molecule or complex and comprising a reagent, the composition being kinetically stable such that gold is deposited on the substrate essentially only where a nucleation center is present thereon, whereby upon such contact, gold atoms from said gold-providing agent are released and deposited onto said nucleation center; and
 - (c) detecting presence of gold on said substrate, such gold signifying presence of said specific separation product on the substrate at a site where the gold is detected.
- 23. A method for assaying presence of an analyte in a sample comprising:
 - (a) providing a substrate carrying capturing agents which bind said analyte;
 - (b) contacting said substrate with said sample, whereby said analyte binds to said capturing agents;
- 15 (c) providing conditions permitting formation of nucleation centers on sites of said substrate comprising said analyte;
- (d) contacting said substrate with a treatment composition comprising a soluble gold-containing molecule or complex and a reagent, the composition being kinetically stable such that gold is deposited on the substrate essentially only at sites
 thereof containing the nucleation centers; and
 - (e) detecting metallic gold deposits on said substrate, indicative of the presence of said analyte in said sample.
 - 24. A method according to Claim 23, wherein step (c) comprises:
- (c1) providing nucleation center-forming agents which comprise each at
 least one moiety, having specific binding affinity to said analyte, coupled to
 at least one nucleation center being one or more of the group consisting of
 metal particle, cluster of metal atoms and a metal-containing complex; and
 - (c2) contacting said substrate with said agents.
- 25. A method according to Claim 24, wherein said analyte is one of a binding couple consisting of an antigen and an antibody or an antibody derivative with an

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antigen-binding domain; sugar and a lectin; a receptor and a ligand; a nucleotide sequence and a complementary nucleotide sequence; a nucleotide sequence and its binding protein or other specific binding agent; biotin and avidin or streptavidin; cellulose or chitin and cellulose binding domain; and said at least one moiety is the other of said binding couple.

- 26. A method according to Claim 23, wherein step (b) comprises:
 - (b1) contacting said sample with nucleation centers, the nucleation centers being one or more of the group consisting of metal particles, cluster of metal atoms and metal-containing complexes, and providing conditions for coupling of said nucleation centers to the analyte if present in the sample, thus obtaining a modified sample containing modified analytes; and
 - (b2) contacting said substrate with said modified analytes bound to said capturing agents.
- 27. A method according to Claim 26, wherein said analyte is one of a binding couple consisting of an antigen and an antibody or an antibody derivative with an antigen-binding domain; sugar and a lectin; a receptor and a ligand; a nucleotide sequence and a complementary nucleotide sequence; a nucleotide sequence and its binding protein or other specific binding agent; biotin and avidin or streptavidin; cellulose or chitin and cellulose binding domain; and said capturing agent is the other of said binding couple.
 - 28. A method according to any one of Claims 23 to 27, wherein said capturing agents are carried on the substrate between electrodes such that gold deposited onto said nucleation centers in step (d) establishes an electric contact between the electrodes; and
 - detection of the gold deposits in step (e) is performed by measuring current-potential relationship between the electrodes.
 - 29. A method according to any one of Claims 23 to 28, wherein said at least one metal particle, cluster of metal atoms or a metal-containing complex is a gold particle or cluster containing gold atoms.

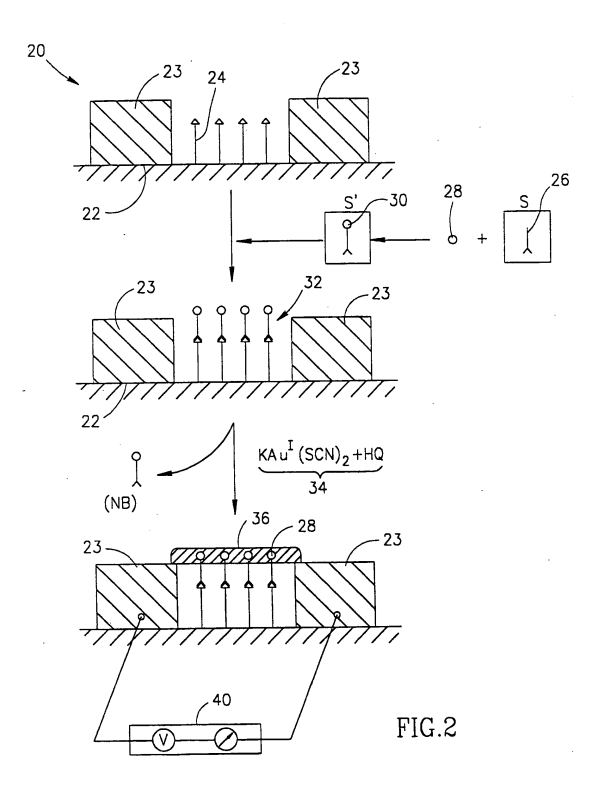
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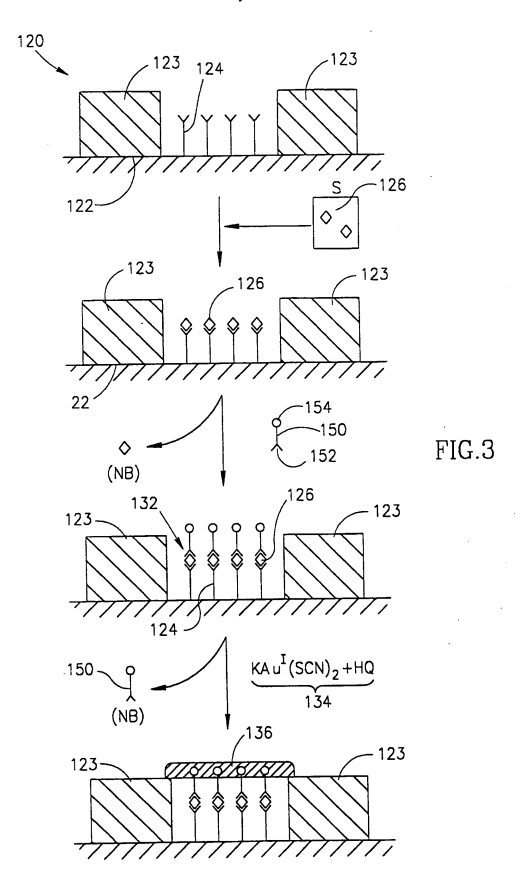
- A method according to any one of Claims 23 to 29, wherein said treatment 30. composition is an aqueous solution.
- A method according to Claim 30, wherein said gold-providing agent is 31. Au^I(SCN)₂.
- A method according to Claim 31, wherein said reagent is a hydroquinone. **32.**
 - A kit for use in the method according to any one of Claims 1 to 32. 33.
- A kit according to Claim 33, comprising a treatment composition which 34. comprises a soluble gold-providing agent, being a gold-containing molecule or complex and comprises a reagent, the composition being kinetically stable such that upon contact with a substrate gold is deposited on a substrate essentially only at sites of the substrate containing nucleation centers.
- A kit according to Claim 33 or 34, further comprising nucleation 35. center-forming agents, for forming nucleation centers at one or more sites on the substrate, said agents comprise each at least one moiety having a specific binding 15 affinity to a substance present at said one or more sites coupled to at least one nucleation center moiety, being one or more of the group consisting of metal particle, cluster of metal atoms and a metal containing complex.
 - A kit for use in a method according to any one of Claims 23 to 32, for 36. assaying an analyte in a sample, comprising:
- a substrate carrying capturing agents which bind said analyte; (i) 20
 - agents for forming nucleation centers at portions of the substrate on (ii) which said analyte becomes immobilized;
- a treatment composition comprising a soluble gold-containing molecule or complex and a reagent, the composition being kinetically stable such 25 that gold is deposited on the substrate essentially only at sites thereof containing the nucleation centers.
 - A kit according to Claim 36, wherein said agent for forming nucleation 37. centers comprises a nucleation center and substances needed in order to couple said nucleation center to said analyte.

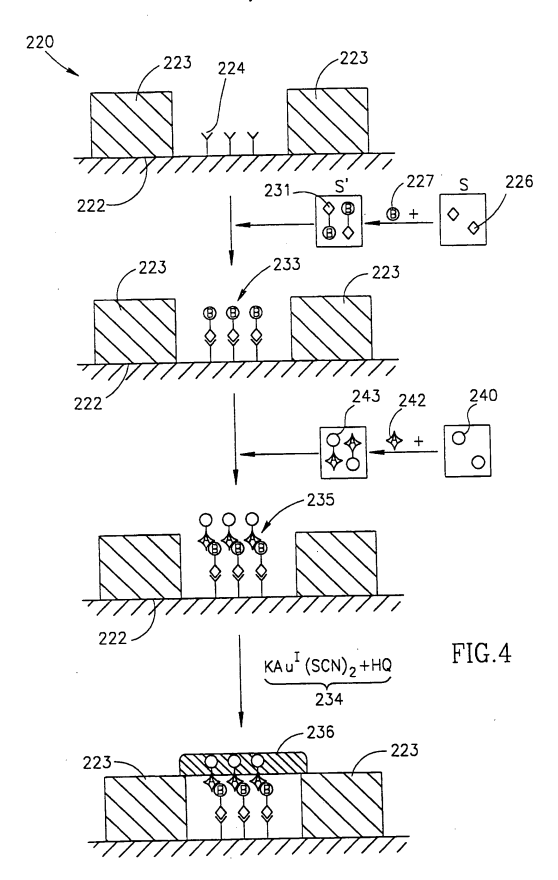
- 38. A kit according to Claim 36, further comprising nucleation center- forming agents which comprise each at least one moiety, having specific binding affinity to said analyte, coupled to at least one nucleation center moiety being one or more of the group consisting of metal particle, cluster of metal atoms and a metal containing complex.
 - 39. A kit according to any one of Claims 36 to 38, wherein said substrate comprises two or more electrodes, and said capturing agents are carried within gaps between the electrodes.

- (a) $KAu^{III}Cl_4$ KSCN KA $u^{III}(SCN)_4$
- (b) KAu (SCN)4 (SPONTANEOUS) KAu (SCN)2
- (c) $HQ + KAu^{I}(SCN)_{2}$ $BQ + Au^{0}$

FIG.1







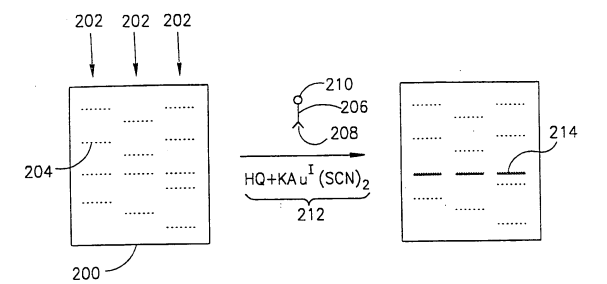


FIG.5

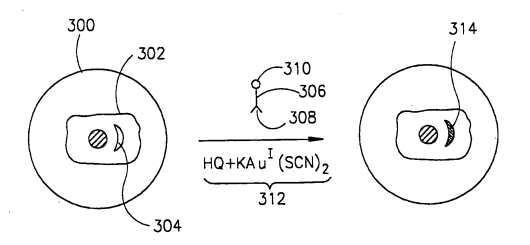
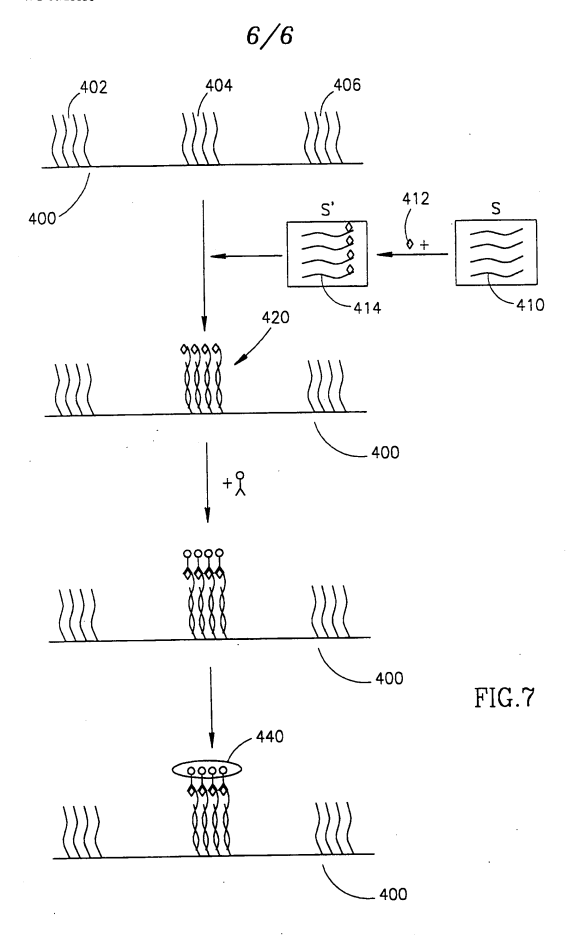


FIG.6



INTERNATIONAL SEARCH REPORT

Inte ional Application No PCT/IL 99/00570

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 G01N33/543 G01N1/30 G01N33/58 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 7 G01N C23C Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Category ° Citation of document, with indication, where appropriate, of the relevant passages 1,6-8,DATABASE WPI Α 16-18, Week 199321 30-32 Derwent Publications Ltd., London, GB; AN 1993-172471 XP002129601 OTAKI HIROAKI: "Spherical fine gold particles prodn. for ceramics or electronic industry" & JP 05 105444 A (MITSUBISHI MATERIALS CORP), 27 April 1993 (1993-04-27) abstract 1,6-8. US 5 202 151 A (USHIO JIRO ET AL) Α 16-18, 13 April 1993 (1993-04-13) 30 - 32column 7, line 57 - line 69; examples 1D.1E -/--Patent family members are listed in annex. Further documents are listed in the continuation of box C. Special categories of cited documents: "T" later document published after the International filling date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed Invention filing date cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another "Y" document of particular relevance; the claimed invertion cannot be considered to involve an inventive step when the document is combined with one or more other such document. citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 14/02/2000 3 February 2000 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tei. (+31-70) 340-2040, Tx. 31 651 epo nl, Hart-Davis, J Fax: (+31-70) 340-3016

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